PATENT SPECIFICATION

(11) 1 581 493

(21) Application No. 21013/78 (31) Convention Application No. 2 722 752

(22) Filed 22 May 1978

(32) Filed 20 May 1977 in

(33) Fed. Rep. of Germany (DE)

(44) Complete Specification published 17 Dec. 1980

(51) INT CL3 COSF 20/12; COSL 25/02, 33/08, 33/10

(52) Index at acceptance

C3P JP C3V BB

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(54) PLASTISOLS

We, RÖHM G.M.B.H., a German Body Corporate of Darmstadt, Germany, do hereby declare the invention for which we pray that a patent may be granted to us, and the method by which it is to be performed, to be particularly described in and by the following statement:

This invention relates to plastisols based on methyl methacrylate copolymers

and organic softeners.

Plastisols are liquid to pasty mixtures which contain a particulate polymer in a liquid organic softener and, as a rule, inorganic fillers and which gel after heating to form solid compositions. Polyvinyl chloride is widely used as the particulate polymer, being particularly suitable for the preparation of plastisols in that it does not dissolve appreciably in the liquid softener at room temperature even after a lengthy storage time. Heating to at least 80 to 100°C results in dissolution of the polyvinyl chloride in the softener, the resulting solution having the properties of a solid owing to the high polymer concentration. After cooling to room temperature

However, polyvinyl chloride does have certain disadvantages as the polymer component of plastisols. Coverings prepared therefrom become yellowed, under the influence of light. Upon heating polyvinyl chloride may liberate hydrogen chloride which can lead to corrosion in production processes where heat is applied. Moreover, hydrogen can occur in dangerously high concentrations in the event of a fire or in the burning of waste. It has therefore been proposed in German Auslegesschrift 2,454,235 to prepare plastisols using methyl methacrylate polymers

and organic softeners.

and organic softeners.

Although homopolymers and copolymers of methyl methacrylate do not have the above-mentioned disadvantages of polyvinyl chloride, they do not possess its special advantages in the gelling operation. Polymethyl methacrylate and any copolymers of methyl methacrylate form together with organic softeners durable plastisols which gel upon heating to e.g. 150°C. However, after the gelled material has been cooled to room temperature, the mixture proves in many cases to be unstable and exudes the softener again in liquid form. This is especially true with those softeners which are used on account of their low price, such as for example, which acid esters. It is possible to increase the compatibility of methyl phthalic acid esters. It is possible to increase the compatibility of methyl methacrylate copolymers with phthalate softeners by using acrylic or methacrylic acid esters of higher alcohols as comonomers. Although the softener-containing gelled masses which are prepared with these copolymers remain homogeneous at room temperature and below over a length of time, the plastisols prepared in this way gel even at room temperature within a few days. Plastisols stable during storage may, in general use, only be obtained with these copolymers if special softeners such as e.g. the relatively expensive dipropylene glycol diberzoate or tricthylhexyl mellitate are used. It is true that to a small extent the storability can also be improved by the use of relatively coarse polymer particles, but this gain in storability is offset by the disadvantage of there being a longer time required for the gelling operation and poorer film properties.

It is an object of the present invention to provide new and advantageous plastisols based on methyl methacrylate polymers.

According to the present invention we provide plastisols comprising an organic softener and an emulsion copolymer in the form of particles having a core/shell construction and consisting of:

	1,301,433	2
5	 (a) a core material compatible with the said softener and comprising a polymer derived from a monomer or monomer composition comprising:— (A) 15 to 100% by weight of at least one monomer selected from alkyl acrylates containing at least 3 carbon atoms in the alkyl moiety, alkyl methacrylates containing at least 2 carbon atoms in the alkyl moiety, and styrene; (B) 0 to 85% by weight of at least one monomer selected from methyl acrylate, methyl methacrylate and ethyl acrylate; and/or (C) 0 to 20% by weight of one or more further radically polymerisable monomers; and 	5
10	(b) a shell material which is incompatible with the said softener, the said shell material comprising a homopolymer of methyl methacrylate or a copolymer containing at least 80% by weight of units of methyl methacrylate and begins a place	10
15	(b) being present in a weight ratio of 3:1 to 1:3, and the said emulsion copolymer and the said organic softener being present in a weight ratio of 10:3 to 1:10. We have prepared plastisols in accordance with the present invention and	15
20	have found that such plastisols are stable for a long period of time even when inexpensive softeners are employed. Such softeners are not exuded at room temperature after gelling. The plastisols according to the invention may, if desired, further contain conventional fillers.	20
25	The core material in the emulsion copolymer particles preferably consists of a copolymer which is distinguished by good compatibility with softeners, particularly with phthalate softeners such as e.g. dioctyl phthalate. The core material alone would gel with the softeners even at room temperature within a short time. A homopolymer or copolymer of methyl methacrylate serves as the shell material, which is stable to softeners at room temperature, but which would not retain the	25 °
30	softener in the homogeneous phase after gelling and cooling. In storage the shell material provides sufficient protection against premature gelling of the core material. If gelling has taken place at high temperature, the affinity of the softener for the core material is such that incompatibility with the shell material no longer leads to separation.	30
35	The compatibility of the core material with softeners is based on its content of alkyl acrylates or methacrylates containing respectively at least three or at least two carbon atoms in the alkyl moiety, and/or its content of styrene. Apart from a few exceptions such as, for example, polyethyl methacrylate, the affinity for the softener, especially a phthalic acid ester softener, of a homopolymer of these	35
40	any permanent protection against gelling at room temperature unless this affinity were attenuated by the presence of a more strongly polar monomer (B). Stability in storage can be considered as sufficient for industrial application if the plastisol does not gel within 21 days at 30°C. A shorter storability of a 2.3 to 5.	40
45	days is sufficient in many cases, for example, when the plastisol is prepared and processed in the same factory. In comparison with plastisols consisting of polymers which are not made in shell form and which often gel within a few hours at room temperature and are therefore of limited industrial use, the plastisols according to the invention represent an important technical advance even with a storability of only a few days because the minimum storage time for industrial usage is still exceeded.	45
50	Since the affinity between the core material and softener depends closely on their chemical composition, the amount of polar monomer (B) is governed by the type of softener, by the type of monomer or the type of type of the type of type of type of the type of type	50
55	shell material. The polymer composition is therefore always adapted to the softener to be used and allows only small variations from the optimum determined at any given time within the limits specified for component (A) and (B). In a preferred case, the core material contains as component (A) and (B). In a preferred	55
50	or methyl methacrylate or mixtures thereof. Besides the monomers (A) and (B), further radically polymerisable monomers (C) can be employed in a proportion up to 20% by weight in the composition of the core material, particularly if their presence is desirable for special response.	60
5	example acrylic and methacrylic acid, their amides, nitriles, hydroxyalkyl esters and amino alkyl esters, vinyl esters of aliphatic carboxylic acids, vinyl pyrrolidone and vinyl imidazole.	65

Polymethyl methacrylate is particularly suitable as shell material. If a copolymer of methyl methacrylate is used instead, the proportion of comonomer or copolymer of methyl methacrylate is used instead, the proportion of comonomer or comonomers should be the smaller, the less polar they are. Monomers of the same type as group (A) are examples of non-polar monomers; their proportion in shell material preferably does not exceed 10%. More strongly polar monomers such as e.g. those of group (B) can form up to 20% by weight of the shell material. The choice of type and quantity of any comonomers optionally used in addition to methyl methacrylate is made such that the shell material, if it is homopolymerised, has a glass temperature of at least 50°C. Small proportions of strongly comonomers in the shell material, e.g. 0.5 to 8% of acrylic or methacrylic acid or acrylamide, have a favourable effect on the storage stability of the plastisols.

Since the plastisols are frequently used as coating agents, especially for metals, 5 5 10 have a favourable effect on the storage stability of the plastisols.

Since the plastisols are frequently used as coating agents, especially for metals, adhesion-assisting monomers are preferably employed in the preparation of the polymer. Examples of such monomers include polymerisable unsaturated carboxylic acids such as acrylic or methacrylic acid, and hydroxyalkyl esters or aminoalkyl esters of these acids. N-vinyl imidazole is especially effective. The adhesion-assisting effect of these comonomers is generally manifested at proportions of 0.1 to 5, preferably 1 to 2%. The adhesion-assisting comonomers can be polymerised into the core material and/or into the shell material.

The importance of the core/shell construction of the emulsion polymer is 10 15 15 20 The importance of the core/shell construction of the emulsion polymer is revealed by a comparison of plastisol prepared according to the invention with 20 those mixtures in which the same softener is used with the same polymer quantity of the core material alone or of the shell material alone or with a mixture of these polymers. A copolymer was used for further comparison, whose overall composition coincided with the polymer composition of the plastisol according to the invention, although it was not prepared in a core-shell form. 25 25

Plastisol consisting of 2 parts of polymer and 3 parts of dioctyl phthalate, storage at 30°C, gelling 30 minutes at 150°C.

Polymer	Stability in storage (in days at 30°C)	Gelled product
Core material 30% BA 70% MMA l:1	>21	Compatible
Copolymer 30% BA, 70% MMA	. 0	Compatible
PMMA	>21	Incompatible
Mixture PMMA + Copolymer 30% BA, 70% MMA (1:1)	0	Compatible
Copolymer 15% BA, 85% MMA	5	Incompatible

BA = n-butyl acrylate, MMA = methyl methacrylate, PMMA = Polymethyl methacrylate.

Determination of the optimal polymer composition for the dioctyl phthalate used as softener is illustrated in Table II.

			IABLE		
		Pol	Polymer		
Example	J	Core	Shell	(in days at 30°C)	Gelled Product
1	20% BA,	80% MMA	MMA	>21	Incompatible
	30% BA,	70% MMA	MMA	>21	Compatible
m	35% BA,	65% MMA	MMA	12	Compatible
4	40% BA,	60% MMA	MMA	e	Compatible
Comparison Exp. 4a	Copolymer: 20% BA,	80% MMA		-	Incompatible
Comparison Exp. 4b	Mixture: 60% MMA 40% BA, 60% MMA + PMMA=1/1	. 60% MMA A=1/1		Solidifies immediately	Compatible
٧	45% BA,	55% MMA	MMA		Compatible
9	50% BA,	50% MMA	MMA	1/2	Compatible
. 7	35% BA,	65% MMA	95% MMA, 5% MAA	17.	Compatible
60	35% BA,	65% MMA	95% MMA, 5% MMAA	, z .	Compatible
6	35% BA,	65% MMA	95% MMA, 5% AS	17.	Compatible
10	35% BA,	65% MMA	95% MMA, 5% MAS.	×211	Compatible
11	40% BA,	60% MMA	80% MMA, 20% MA	e	Compatible
. 21	15% nBMA,	85% MMA	MMA	>21	Incompatible
13	50% nBMA, 50% MMA	50% MMA	MMA	>21	Compatible
Comparison Exp. 13a Copolymer: 25% nBMA, 75% MMA	Copolymer: 25% nBMA,	75% MMA		1 2<	Compatible
Comparison Exp. 13b	Mixture: 50% nBMA, 50% NMA +MMA-1/1	50% MMA A=1/1		Solidifies immediately	Compatible

TABLE

			TABLE (Continued)		
		Po	Polymer		
Example		Core	Shell	(in days at 30°C)	Gelled Product
14	70% nBMA, 30% MMA	30% MMA	MMA	9	Compatible
15	70% nBMA, 5% MAS	70% nBMA, 25% MMA, 5% MAS	MMA	>21	Compatible
16	70% nBMA, 30% MA	30% MA	MMA	m	Compatible
. 11	20% nBma, 30% St	50% MMA,	MMA	. >21	Compatible
. 18	10% EHA,	90% MMA	MMA	>21	Incompatible
19	20% EHA,	80% MMA	MMA	>21	Compatible
20	30% EHA,	70% MMA	MMA	19	Comparible
11	100% EMA		MMA	>21	Compatible
22	10% EMA,	30% MMA .	MMA	×21	Compatible
23	80% i.BMA, 20% MMA	20% MMA	MMA	>21	Compatible
24	20% EHMA, 80% MMA	80% MMA	MMA	>21	Compatible
25	30% BA, 1% VI	69% MMA,	99% MMA, 1% VJ	>21	Compatible
Explanation of the attention					

Methyl acrylate
Methyl methacrylate
Ethyl methacrylate
n-butylacrylate
A n-butylmethacrylate
methacrylic acid amide

Isobutyl methacrylate
Ethylhexyl methacrylate
Ethylhexyl acrylate
Acrylic acid
Methacrylic acid
Styrene i.BMA EHMA EHA AS MAS St

	1,381,493	8
	Whereas in Examples 1 to 25 the ratio by weight of core material to s material is 1:1, in Examples 26 and 27 it is adjusted to 2:1 and 1:2 respective	hell ely.
5	Example 26 In a Witt pot (2 litres) with reflux cooler, agitator and feed vessel 0.1 ammonium peroxide persulphate and 3 g of C ₁₈ -paraffin sulphonate (trade na emulsifier K30 Bayer AG) are dissolved at 80°C in 400 g of distilled was Emulsion 1 is added to this solution with stirring over 2 hours at 80°C.	g of me: 5 iter.
	Emulsion I (core material): 417 g of MMA	
	250 g of BA	
10	3.13 g of emulsifier	10
	0.20 g of initiator	
	400 g of distilled water.	
	Emulsion 2 is subsequently added over 1 hour	
	Emulsion 2 (shell material): 333 g of MMA	
15	1.57 g of emulsifier	15
	0.10 g of initiator	
	200 g of distilled water.	
20	After everything has been added, the mixture is kept at 80°C for 2 hours at then cooled to approximately 25°C and spray-dried. The plastisol properties tested as in Examples 1 to 25.	and are 20
	Stability in storage at 30°C: >21 days.	
	Gelled product: compatible.	
25	Example 27 The procedure is as in Example 26 with the difference that the follower emulsion is added over 1 hour:	ing 25
	Emulsion 1 (core material): 250 g of nBMA	
	83 g of MMA	
	1.57 g of emulsifier	
	0.1 g of initiator	
Ю	200 g of distilled water.	30
	Emulsion 2 is subsequently added over 2 hours.	
	Emulsion 2 (shell material): 666 g of MMA	
	3.13 g of emulsifier	
	0.2 g of initiator	
5	400 g of distilled water.	35
	After everything is added the mixture is kept at 80°C for 2 hours and the	eņ

After everything is added the mixture is kept at 80°C for 2 hours and then cooled to 25°C and spray-dried. The plastisol properites are tested as in Examples 1 to 25.

Degreased iron sheets are coated by dipping and stoved for 10 minutes at 180°C. Strongly adhering, flexible coverings are obtained in all cases.

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30 parts of chalk

10 parts of titanium dioxide.

	Example 31 A kneadable filler mass is prepared from:	
	20 parts of polymer powder according to Example 25	
	30 parts of dioctyl phthalate	
5	60 parts of chalk	5
	I part of micronised porous silicic acid.	
	The kneadable mass is applied by hand to a thickness of approximately 5 mm on an electrophoretically primed sheet and stoved for 25 minutes at 120°C. A strongly adhering, flexible covering is obtained.	
10	WHAT WE CLAIM IS:— 1. Plastisols comprising an organic softener and an emulsion copolymer in the form of particles having a core/shell construction and consisting of:— (a) a core material compatible with the said softener and comprising a polymer desired from	10
15	derived from a monomer or monomer composition comprising: (A) 15 to 100% by weight of at least one monomer selected from alkyl acrylates containing at least 3 carbon atoms in the alkyl moiety, alkyl methacrylates containing at least 2 carbon atoms in the alkyl moiety, and styrene; (B) 0 to 85% by weight of at least one monomer selected from methyl acrylate,	15
20	(C) 0 to 20% by weight of one or more further radically polymerisable monomers; and	. 20
25	(b) a shell material which is incompatible with the said softener, the said shell material comprising a homopolymer of methyl methacrylate or a copolymer containing at least 80% by weight of units of methyl methacrylate and having a glass temperature of at least 50°C; the said core material (a) and the said shell material (b) being present in a weight ratio of 3:1 to 1:3, and the said emulsion copolymer	25
30	and the said organic softener being present in a weight ratio of 10:3 to 1:10. 2. Plastisols as claimed in claim 1 wherein component (A) of the said monomer composition comprises an alkyl methacrylate containing 4 to 18 carbon atoms in the alkyl moiety. 3. Plastisols as claimed in claim 1 or claim 2 wherein component (C) of the said	30
35	monomer composition comprises at least one monomer selected from acrylic and methacrylic acid; amides, nitriles, hydroxyalkyl esters and aminoalkyl esters of such acids; vinyl esters of aliphatic carboxylic acids; vinyl pyrrolidone; and vinyl imidazole. 4. Plastisols as claimed in any of the preceding claims wherein the said core	35
40	material and/or the said shell material contain(s) units of at least one adhesion assisting monomer selected from polymerisable unsaturated carboxylic acids and hydroxyalkyl and aminoalkyl esters of such acids; and N-vinyl-imidazole. 5. Plastisols as claimed in any of the preceding claims wherein the said softener comprises an ester of phthalic acid.	40
45	 6. Plastisols as claimed in claim 5 wherein the said softener comprises dioctyl phthalate or dibutyl phthalate. 7. Plastisols as claimed in any of claims 1 to 4 wherein the said softener comprises an ester of sebacic or azelaic acid or a polymeric softener. 8. Plastisols as claimed in any of the preceding claims containing at least one 	45
50	filler. 9. Plastisols as claimed in claim 8 wherein the said filler comprises chalk, kaolin or mica powder. 10. Plastisols as claimed in any of the preceding claims wherein the said emulsion copolymer and the said organic softener are present in a weight ratio of	50
55	2:3 to 1:2. 11. Plastisols as claimed in claim 1 substantially as herein described. 12. Plastisols as claimed in claim 1 substantially as herein described in any of the Examples. 13. A method of coating a substrate which comprises applying a coating of a plastisol as claimed in any of the preceding claims to a substrate and subsequently	55
	gelling the plastisol coating.	

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14. A method as claimed in claim 13 wherein the plastisol coating is applied in

a thickness of 5 μm to 5 mm.

15. A method as claimed in claim 13 wherein the plastisol coating is applied in 15. A method as claimed in claim 13 or claim 14 wherein the plastisol coating is gelled at 90 to 200°C.

16. A method as claimed in claim 13 substantially as herein described.

17. A method as claimed in claim 13 substantially as herein described in any of the Examples.

18. Coated substrates whenever prepared by a method as claimed in any of claims 13 to 17.

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Printed for Her Majesty's Stationery Office by the Courier Press, Learnington Spa. 1980. Published by the Patent Office, 25 Southampton Buildings, London, WC2A 1AY, from which copies may be obtained.